

Tests that diagnose diseases are less reliable than you'd expect. Here's why

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Credit: CDC / Unsplash

You feel unwell, and visit your doctor. They ask some questions and take some blood for testing; a few days later they call to say you have been diagnosed with a disease.

What are the chances you actually have the disease? For some common diagnostic tests, the answer is surprisingly low.



Few <u>medical tests</u> are 100% accurate. Part of the reason is that people are inherently variable, but many tests are also built on limited or biased samples of patients—and our own work has shown researchers may <u>deliberately exaggerate</u> the effectiveness of new tests.

None of this means we should stop trusting diagnostic tests, but a better understanding of their strengths and weaknesses is essential if we want to use them wisely.

People are variable

An example of a widely used imperfect test is <u>prostate-specific antigen</u> (PSA) screening, which measures the level of a particular protein in the blood as an indicator of prostate cancer.

The test catches an estimated 93% of cancers—but it has a very high false positive rate, as around 80% of men with a positive result do not actually have cancer. For those in the 80%, the result creates unnecessary stress and likely further testing including painful biopsies.

Rapid antigen tests for COVID-19 are another widely used imperfect test. A <u>review of these tests</u> found that, of people without symptoms but with a positive test result, only 52% actually had COVID.

Among people with COVID symptoms and a positive result, the accuracy of the tests rose to 89%. This shows how a test's performance cannot be summarized by a single number and depends on individual context.

Why aren't <u>diagnostic tests</u> perfect? One key reason is that people are variable. A high temperature for you, for example, might be perfectly normal for someone else. For blood tests, many extraneous factors can influence the results, such as the time of day or how recently you have



eaten.

Even the ubiquitous blood pressure test <u>can be inaccurate</u>. Results can vary depending on whether the cuff is a good fit for your arm, if you have your legs crossed, and if you're talking when the test is done.

Small samples and statistical skullduggery

There's an enormous amount of research on new diagnostic models. New models frequently make the headlines as "medical breakthroughs," such as how your <u>handwriting could detect Parkinson's disease</u>, how your pharmacy loyalty card could <u>detect ovarian cancer earlier</u>, or how <u>eye</u> <u>movements could detect schizophrenia</u>.

But living up to the headlines is often a different story.

Many diagnostic models are developed based on small sample sizes. <u>A</u> review found half of diagnostic studies used just over 100 patients. It is hard to get a true picture of the accuracy of a diagnostic test from such small samples.

For accurate results, the patients who use the test should be similar to those who were used to develop the test. For example, the widely used Framingham Risk Score for identifying people at high risk of heart disease was developed in the United States and is known to <u>perform</u> <u>poorly</u> in Aboriginal and Torres Strait Islander people.

Similar disparities in accuracy have been found for "polygenic risk scores." These combine information on thousands of genes to predict disease risk, but were developed in European populations and <u>perform</u> poorly in non-European populations.

Recently, we identified another important problem: researchers have



exaggerated the accuracy of some models to gain journal publications.

There are many ways to exaggerate the performance of a test, such as dropping hard-to-predict patients from the sample. Some tests are also not truly predictive, as they include information from the future, such as a <u>predictive model of infection</u> that includes whether the patient had been prescribed antibiotics.

Perhaps the most extreme example of exaggerating the power of a diagnostic test was the Theranos scandal, in which a finger-prick blood test supposed to diagnose multiple health conditions attracted hundreds of millions of dollars from investors. This was too good to be true—and the mastermind has now been convicted of fraud.

Big data can't make tests perfect

In the era of precision medicine and <u>big data</u>, it seems appealing to combine tens or hundreds of pieces of information about a patient—perhaps using machine learning or artificial intelligence—to provide highly accurate predictions. However, the promise is so far outstripping the reality.

One <u>study</u> estimated 80,000 new prediction models were published between 1995 and 2020. That's around 250 new models every month.

Are these models transforming health care? We see no sign of it—and if they really were having a big impact, surely we wouldn't need such a steady stream of new models.

For many diseases there are data problems that no amount of sophisticated modeling can fix, such as measurement errors or missing data that make accurate predictions impossible.



Some diseases or illnesses are likely inherently random, and involve complex chains of events which a patient cannot describe and no <u>model</u> could predict. Examples might include injuries or previous illnesses that happened to a patient decades ago, which they cannot recall and are not in their medical notes.

Diagnostic tests will never be perfect. Acknowledging their imperfections will enable doctors and their patients to have an informed discussion about what a result means—and most importantly, what to do next.

More information: Nicole White et al, Evidence of questionable research practices in clinical prediction models, *BMC Medicine* (2023). DOI: 10.1186/s12916-023-03048-6

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